TENT COOPERATION TRE

	From the INTERNATIONAL BUREAU
PCT	То:
NOTIFICATION OF ELECTION (PCT Rule 61.2)	Assistant Commissioner for Patents United States Patent and Trademark Office Box PCT Washington, D.C.20231 ETATS-UNIS D'AMERIQUE
Date of mailing (day/month/year)	TATS-ONIS D'AMERIQUE
13 October 2000 (13.10.00)	in its capacity as elected Office
International application No. PCT/DK00/00040	Applicant's or agent's file reference 23923PC1
International filing date (day/month/year) 01 February 2000 (01.02.00)	Priority date (day/month/year) 03 February 1999 (03.02.99)
Applicant REES, Stephen, Edward et al	
REES, Stephen, Luward et al	
The designated Office is hereby notified of its election ma in the demand filed with the International Prelimina 07 August 20 in a notice effecting later election filed with the Inte 2. The election X was was not made before the expiration of 19 months from the priority Rule 32.2(b).	ory Examining Authority on:
<u> </u>	
The International Bureau of WIPO 34, chemin des Colombettes 1211 Geneva 20, Switzerland	Authorized officer Beatriz Morariu

Facsimile No.: (41-22) 740.14.35 Form PCT/IB/331 (July 1992)

DK0000040

Telephone No.: (41-22) 338.83.38

PCT/DK00/00040

Copy for the Elected Office (EO/US)

PATENT COOPERATION TRI TY

		From th	ne INTERN	ATIONAL B	UREAU
PCT		To:			
NOTIFICATION OF THE RECORDING OF A CHANGE (PCT Rule 92bis.1 and Administrative Instructions, Section 422)		PLOUGMANN, VINGTOFT & PARTNERS A/S Sankt Annæ Plads 11 P.O. Box 3007 DK-1021 Copenhagen K. DANEMARK			
12 July 2001 (12.07.01)					-
Applicant's or agent's file reference 23923PC1			IMPOR	TANT NOT	IFICATION
International application No. PCT/DK00/00040			-	e (day/month/y 100 (01.02.00	· ·
FC17DR00/00040		011	ebidary 20	00 (01.02.00	· · · · · · · · · · · · · · · · · · ·
The following indications appeared on record conce X the applicant X the inventor	erning:	the ager	it [the comm	on representative
Name and Address			State of Na	tionality	State of Residence
REES, Stephen, Edward Vesterbro 60, 5.th.			GB Telephone	No	DK
DK-9000 Aalborg Denmark			relephone	140.	
- Sommerk			Facsimile N	lo.	
			Teleprinter	No.	
2. The International Bureau hereby notifies the applica		ī	一 ·		
the person the name X	the add	ress	the nati	onality	the residence
Name and Address			State of Na	tionality	State of Residence
REES, Stephen, Edward Forchhammersvej 40 DK-9000 Aalborg			Telephone	No.	
Denmark			Facsimile N	lo.	
			Teleprinter	No.	
3. Further observations, if necessary:					
4. A copy of this notification has been sent to:					
X the receiving Office		[the desi	gnated Offices	concerned
the International Searching Authority		ĺ	X the elec	ted Offices cor	ncerned
X the International Preliminary Examining Author	rity	[other:		
		Authorized	officer		
The International Bureau of WiPO 34, chemin des Colombettes				J. Leitao	
1211 Geneva 20, Switzerland Facsimile No.: (41-22) 740.14.35		Telephone	No.: (41-22)	338.83.38	

Form PCT/IB/306 (March 1994)



INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference	FOR FURTHER see Notification (Form PCT/ISA/	of Transmittal of International Search Report 220) as well as, where applicable, item 5 below.				
23923PC1	ACTION					
International application No.	International filing date (day/month/year) (Earliest) Priority Date (day/month/year)					
PCT/DK 00/00040 01/02/2000 03/02/1999						
Applicant						
REES, Stephen, Edward et	al.					
This International Search Report has bee according to Article 18. A copy is being to	en prepared by this International Searching Aut ransmitted to the International Bureau.	hority and is transmitted to the applicant				
	2					
This International Search Report consists It is also accompanied b	s of a total of Sheets. y a copy of each prior art document cited in this	report.				
it is also accompanied o	y a copy of each pilot air decament executive					
1. Basis of the report						
With regard to the language, the language in which it was filed, ur	e international search was carried out on the ba nless otherwise indicated under this item.	sis of the international application in the				
the international search Authority (Rule 23.1(b)).	was carried out on the basis of a translation of	the international application furnished to this				
b With regard to any nucleotide a	nd/or amino acid sequence disclosed in the i	nternational application, the international search				
was carried out on the basis of the	he sequence listing: ional application in written form.					
·	ternational application in computer readable for	m.				
·	to this Authority in written form.					
· —	to this Authority in computer readble form.					
	ubsequently furnished written sequence listing o	does not go beyond the disclosure in the				
international application	as filed has been furnished.	•				
the statement that the in furnished	formation recorded in computer readable form	is identical to the written sequence listing has been				
. —	und unsearchable (See Box I).					
3. Unity of invention is la	cking (see Box II).					
A NACAL						
4. With regard to the title , X the text is approved as s	submitted by the applicant.					
	ished by this Authority to read as follows:					
Life text has been established by this reading to read as renewe.						
·						
5. With regard to the abstract,						
X the text is approved as a	submitted by the applicant.					
the text has been estable within one month from the	lished, according to Rule 38.2(b), by this Author he date of mailing of this international search re	ity as it appears in Box III. The applicant may,. port, submit comments to this Authority.				
6. The figure of the drawings to be pu	blished with the abstract is Figure No.	<u>5</u>				
X as suggested by the app	plicant.	None of the figures.				
because the applicant for	ailed to suggest a figure.					
because this figure better characterizes the invention.						

A. CLASS	IFICATION OF SUBJECT MATTER			
IPC7: A61B 5/08, A61M 16/00 According to International Patent Classification (IPC) or to both national classification and IPC				
	S SEARCHED cumentation searched (classification system followed by	classification symbols)		
		,		
IPC7: A	61B, A61M		the fields searched	
Documentat	ion searched other than minimum documentation to the	extent that such documents are included in	the helds sea ones	
Electronic da	ata base consulted during the international search (name	of data base and, where practicable, search	terms usea)	
C. DOCU	MENTS CONSIDERED TO BE RELEVANT			
Category*	Citation of document, with indication, where app	ropriate, of the relevant passages	Relevant to claim No.	
X EP 0753320 A1 (B. LACHMANN), 15 January 1997 (15.01.97), column 7, line 41 - column 9, line 2, abstract			1-3,31	
A	column 7, line 41 - column 9, line 2, abstract 4-30,32-51			
A	US 5103814 A (T. MAHER), 14 April 1992 (14.04.92), column 3, line 3 - line 55			
A	EP 0502270 A1 (HAMAMATSU PHOTONICS K.K.), 9 Sept 1992 (09.09.92), page 4, line 46 - page 5, line 4, abstract			
			·	
Furth	er documents are listed in the continuation of Box	See patent family anne.	X	
"A" docum	categories of cited documents: ent defining the general state of the art which is not considered	"T" later document published after the int date and not in conflict with the appli the principle or theory underlying the	cation but cited to understand	
"E" erlier d	f particular relevance locument but published on or after the international filing date ent which may throw doubts on priority claim(s) or which is	"X" document of particular relevance: the considered novel or cannot be considered when the document is taken alon	ered to involve an inventive	
cited to establish the publication date of another citation or other special reason (as specified) "O" document referring to an oral disclosure, use, exhibition or other means "O" an oral disclosure, use, exhibition or other means				
the pri	"P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family			
Date of th	Date of the actual completion of the international search Date of mailing of the international search report			
25 May	25 May 2000 25 May 2000			
Name and mailing address of the International Searching Authority European Patent Office P.B. 5818 Patentlaan 2 NL-2280 HV Rijswijk Ulrika Andersson/AE				
Tel(+31-70)340-2040, Tx 31 651 epo nt, Fax(+31-70)340-3016 Telephone No. Telephone No.				

INTERNATION A Information on pa EARCH REPORT a family members

ional application No. 02/12/99 | PCT/DK 00/00040

SA 267044 -

	atent document I in search repor	·t	Publication date		Patent family member(s)	Publication date
EP	0753320	A1	15/01/97	JP SE US	9024099 A 9502543 D 5752509 A	28/01/97 00/00/00 19/05/98
US	5103814	Α	14/04/92	NONI		
EP	0502270	A1	09/09/92	DE JP US	69123954 D,T 5084233 A 5251632 A	30/04/97 06/04/93 12/10/93

PATENT 0459-0638P

IN THE U.S. PATENT AND TRADEMARK OFFICE

Applicant:

REES, Stephen Edward et al.

Int'l. Appl. No.:

PCT/DK00/00040

Appl. No.:

New

Group:

Filed:

August 3, 2001

Examiner:

For:

AUTOMATIC LUNG PARAMETER ESTIMATOR

LETTER

BOX PATENT APPLICATION

Assistant Commissioner for Patents Washington, D.C. 20231

August 3, 2001

Sir:

The PTO is requested to use the amended sheets/claims attached hereto (which correspond to Article 34 amendments or to claims attached to the International Preliminary Examination Report) during prosecution of the above-identified national phase PCT application.

Respectfully submitted,

BIRCH, STEWART, KOLASCH & BIRCH, LLP

By

John A. Castellano, #35,094

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JAC/cqc

0459-0638P

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(703) 205-8000



PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference	FOR FURTHER ACTION		n of Transmittal of International amination Report (Form PCT/IPEA/416)		
International application No. International filing date (onth year) Pr	iority date (day/month/year)		
PCT/DK00/00040	01.02.2000	0	3.02.1999		
International Patent Classification (IPC) o	r national classification and IPC	7			
A 61 B 5/08, A 61 M 1	6/00				
		•			
Applicant					
REESE, Stephan Edward	et al				
been amended and are the t	e applicant according to Article of 4 sheets, inclusioned by ANNEXES, i.e., sheets	36. ding this cover she of the description, containing rectific	eet. claims and/or drawings which have cations made before this Authority		
These annexes consist of a total of	of 11 sheets.				
3. This report contains indications relating to the following items:					
I 🔀 Basis of the report	I Basis of the report				
II Priority					
	f opinion with regard to novelty,	:	Lindonesia Landina (1964)		
		inventive step and	i muusutai appiicaomty		
IV Lack of unity of inve	ention		. •		
citations and explana	ations supporting such statement	o novelty, inventiv	e step or industrial applicability;		
VI Certain documents c	ited				
VII Certain defects in the	e international application				
VIII Certain observations	on the international application				
Date of submission of the demand	Date	of completion of t	his report		
07.08.2000	12	06.2001			
Name and mailing address of the IPEA/S		orized officer	.		
Box 5055	Box 5055 Telex				
S-102 42 STOCKHOLM PATOREG-S Rune Bengtsson/EE					
Facsimile No. 08-667 72 88 Telephone No. 08-782 25 00					

International application No.

PCT/DK00/00040

I.	Bas	of the report
1	. With	egard to the elements of the international application:*
		the international application as originally filed
	\boxtimes	the description:
		pages 1-25 , as originally filed
		pages, filed with the demand
		pages, filed with the letter of
	\boxtimes	the claims:
		pages, as originally filed
		pages, as amended (together with any statement) under article 19
		pages, filed with the demand
	K 7	pages 1-11 , filed with the letter of 27.04.2001
	\bowtie	the drawings:
		pages $1-10$, as originally filed
		pages, filed with the demand
		pages, filed with the letter of the sequence listing part of the description:
	Ш.	
		,
		pages, filed with the demand pages, filed with the letter of
3	These	ernational application was filed, unless otherwise indicated under this item. elements were available or furnished to this Authority in the following language which is: the language of a translation furnished for the purposes of international search (under Rule 23.1(b)). the language of publication of the international application (under Rule 48.3(b)). the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3). egard to any nucleotide and/or amino acid sequence disclosed in the international application, the international inary examination was carried out on the basis of the sequence listing: contained in the international application in written form. filed together with the international application in computer readable form. furnished subsequently to this Authority in written form. furnished subsequently to this Authority in computer readable form. The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the
		international application as filed has been furnished. The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.
4	1.	The amendments have resulted in the cancellation of:
		the description, pages
		the claims, Nos.
		the drawings, sheet/fig
. 5	5.	This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2 (c)).**
*	in th	cement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to s report as "originally filed" and are annexed to this report since they do not contain amendments (Rules 70.16 0.17).
**	Any	eplacement sheet containing such amendments must be referred to under item I and annexed to this report.

International application No.

PCT/DK00/00040

V.	Reasoned statement under Article 35(2) with regard to	novelty, inventive step or industrial applicability;
	citations and explanations supporting such statement	•

1. Statement

Novelty (N)	Claims Claims	1-50	YES NO
Inventive step (IS)	Claims Claims	1-50	YES NO.
Industrial applicability (IA)	Claims Claims	1-50	YES

2. Citations and explanations (Rule 70.7)

The claimed invention relates to an automatic lung parameter estimator, which has a gas-mixing unit supplying a first homogeneous gas to the inlet of a ventilator via a gas mixer. A second gas is mixed with the first gas in the mixer. A computer determines respiratory parameters and controls the gas mixture supplied to the patient.

The following relevant documents were cited in the International Search Report:

D1: EP 0753320, A1 D2: US 5103814, A D3: EP 0502270, A1

Document D1, which is the closest prior art, describes artificial ventilation system. The system comprises respiratory gas delivery a unit with regulatory connectable to the lung of the patient and a parametermonitoring unit comprising a blood gas analyser.

The documents D2 and D3 are two types of respirator system, which both differ significantly from the invention.

The cited document D2 relates to a self-compensating patient respirator. The ventilator comprises a respirator and pulseoximeter for determining the body oxygen saturation of a patient. An oxygen comparison circuit compares the body oxygen saturation to a predetermined body oxygen saturation level. A control is coupled to the respirator, for periodically. decreasing the percentage of oxygen in a respiratory gas within the ventilator while the body oxygen saturation is greater than the predetermined body oxygen saturation level.

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International application No.

PCT/DK00/00040

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation of: V

Document D3 relates to tissue oxygen measuring system, which has a ventilation unit, a control unit for controlling the gaseous content of the air and an arrangement for producing trigger signals. A measurement system is included for measuring oxygen in the tissue and supplying data regarding the measured results, and a data processing unit computing information regarding blood flowing in the tissue.

From D1device a is known for determining respiratory parameters. The parameters that are determined in D1 are either settings for the artificial ventilation system or functions of the measured parameters over time. No parameters in D1 being descriptive of the pulmonary gas exchange are computed as the purpose of the artificial ventilation system. Disclosed is to obtain an optimised artificial ventilation of a lung system while preventing injuries of the lungs. invention according to claims 1-50 is therefore new and involves an inventive step.

The invention according to claims 1-50 is novel (N) and is considered to involve an inventive step (IS). The invention according to claims 1-50 is considered to have industrial applicability (IA).

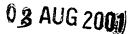
CLAIMS

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1. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FE'O2, FEO2,

- 20 PIO2, PEO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly, the computer being adapted for retrieving and storing at least two measurements being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the at least two measurements being conducted at respective levels of oxygen in the gas flow passing into the respiratory system, the computer further being adapted for determining at least one respiratory parameter (Rdiff, shunt, \(\vert \cdot \vert \ver
 - 2. A device according to claim 1, wherein the computer further being adapted for determining at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual.

- 3. A device according to claim 1 or 2, wherein said parameter(s) (Rdiff, shunt, V/Q, H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 4. A device according to claim 1 or 3, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising

determining, based on at least two measurements, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection 10 means,

producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

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5. A device according to any of claims 1-4, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FEO2, PE'O2, PEO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second

detection means retrieved simultaneously.

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6. A device according to claim 5, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.

- 7. A. device according to any of claims 1-6, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item
 5 accordingly if said parameter exceeds said threshold value.
- 8. A device according to any of claim 1-7, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2,
 10 SpO2, PaO2, PpO2) and produce a control data item accordingly.
- 9. A device according to claim 8, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurements.
 - 10. A device according to claim 8, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 11. A device according to any of claims 8-10, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 12. A device according to any of claims 1-11, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.
 - 13. A device according to any of claims 1-12, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 14. A device according to any of the preceding claims, wherein the oxygen saturation in35 the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.

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15. A device according to any of claims 1-14, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

16. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2)

20 in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FE'O2, FEO2,

PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

25 the computer being adapted for retrieving and storing a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer being further adapted for performing a procedure at least once, the procedure comprising

determining, based on data stored within the data structure, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection means,

producing a possible control data item based on the target, and

retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

5 17. A device according to claim 16, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure in data storage means associated with the computer, in which the stored outputs are mutually related and related to the output from the first detection means and the second detection means, and the output from the four detection means can be retrieved simultaneously.
 - 18. A device according to claim 17, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 19. A device according to claim 16 or 17, wherein the computer is adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 30 20. A device according to claim 19, wherein at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.

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- 21. A device according to claim 19 or 20, wherein said parameter(s) (Rdiff, shunt, \vec{V} / \vec{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 22. A. device according to any of claims 16-21, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.
 - 23. A device according to any of claims 16-22, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.
 - 24. A device according to claim 23, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).
 - 25. A device according to claim 23, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 25 26. A device according to any of claims 23-25, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 27. A device according to any of claims 16-26, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

- 28. A device according to any of claims 16-27, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 29. A device according to any of claims 16-29, wherein the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
 - 30. A device according to any of claims 16-29, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

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- 31. A device for determining one or more respiratory parameters relating to an individual, comprising
- a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,
 - a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2)
25 in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2, FEO2, PIO2, PE'O2, FEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing at least a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer further being adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from

the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

- 32. A device according to claim 31, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).
- 33. A device according to claim 31, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 34. A device according to any of claims 31-33, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one
 15 gas, in response to said control data item from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 35. A device according to any of claims 31 to 34, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising
- determining, based on at least one measurement, whether additional measurements are required,
 - asserting a possible desired target defining a desired output of the first detection means,
- producing a possible control data item based on the target, and
 retrieving and storing, in the data structure, additional measurement results being
 the concurrent output produced by the first detection means and the second detection
 means.
- 36. A device according to any of claims 31-35, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, V) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously.
- 10 37. A device according to claim 36, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 38. A device according to any of claims 31-37, wherein the computer is adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 39. A device according to claim 38, wherein at least two respiratory parameters (Rdiff, 20 shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.
 - 40. A device according to claim 38 or 39, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 41. A. device according to any of claims 31-40, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.
 - 42. A device according to any of claims 31-41, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

- 43. A device according to any of claims 31-42, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 5 44. A device according to any of claims 31-43, wherein the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
- 45. A device according to any of claims 31-43, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.
 - 46. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is an apparently healthy individual.
 - 47. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is considered to have a risk of suffering from hypoxemia.
- 20 48. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is suffering from hypoxemia.
- 49. Method according to claim 48, wherein the individual is suffering from one or more disease(s) selected from the group(s) comprising left sided heart failure, adult respiratory distress syndrome, pneumonia, postoperative hypoxemia, pulmonary fibrosis, toxic pulmonary lymphoedema, pulmonary embolisms, chronic obstructive pulmonary disease and cardiac shunting.
- 30 50. A computer system comprising at least one general purpose computer having one or more computer programs stored within data storage means associated therewith, the computer system being arranged for as well as being adapted for determining one or more respiratory parameters according to any of claims 1-49.

51. A computer program product being adapted to enable a computer system comprising at least one general purpose computer having data storage means associated therewith and being arranged suitably to determine one or more respiratory parameters according to any of claims 1-49.

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PTO/PGT Rec'd 03 AUG	1200 and Application No.
REQUEST	
REQUEST	International Filing Date
The undersigned requests that the present international application be processed	
according to the Patent Cooperation Treaty.	Name of receiving Office and "PCT International Application"
	Applicant s or agent s file reference (if desired) (12 characters maximum) 23923 PC1
Box No. 1 TITLE OF INVENTION	<u></u>
AUTOMATIC LUNG PARAMETER ESTIMATOR	\checkmark
Box No: II APPLICANT	
Name and address: (Family name followed by given name; for a	lacal antity full official
designation. The address must include postal code and name of co address indicated in this Box is the applicant's State (that is, country of residence is indicated below.)	untry. The country of the This person is also inventor
REESE, Stephen Edward	Telephone No.
Vesterbro 60, 5. th.	Facsimile No.
DK-9000 Aalborg Denmark	
Delinar	Teleprinter No.
State (that is, country) of nationality:	State (that is, country) of residence:
GB V	DK V
	ed States except the United States the States indicated in States of America of America only the Supplemental Box
Box No. III FURTHER APPLICANT(S) AND/OR (FURT	THER) INVENTOR(S)
Name and address: (Family name followed by given name; for a designation. The address must include postal code and name of co	untry. The country of the This person is:
address indicated in this Box is the applicant's State (that is, country of residence is indicated below.)	y) of residence if no State applicant only
TOFT, Egon Steen	
Blegdalsparken 102	applicant and inventor
DK-9000 Aalborg Denmark	inventor only (If this check-box
Somman	is marked, do not fill in below.)
State (that is, country) of nationality:	State (that is, country) of residence:
DK	DK O
This person is applicant for the purposes of States all designated the United	ed States except States of America the United States the States indicated in the Supplemental Box
Further populicants and/or (further) inventors are indicated	on a continuation sheet.
Box No. IV AGENT OR COMMON REPRESENTATIV	E; OR ADDRESS FOR CORRESPONDENCE
The person identified below is hereby/has been appointed to act of the applicant(s) before the competent International Authoritie	s as:
Name and address: (Family name followed by given name; for designation. The address must include postal	code and name of country.)
Plougmann, Vingtoft & Partners A/S	+ 45 33 63 93 00
Sankt Annæ Plads 11	Facsimile No. + 45 33 63 96 00
P.O. Box 3007 DK-1021 Copenhagen K.	',
Denmark	Teleprinter No.
Address for correspondence: Mark this check-hov where	no agent or common representative is/has been appointed and the
space above is used instead to indicate a special address to	which correspondence should be sent.
Form PCT/RO/101 (first sheet) (July 1998; reprint January 2000)	See Notes to the request form

Continuation of Box No. III FURTHER APPLICANT(S) AND/OR (FURTHER) INVENTOR(S)			
If none of the following sub-boxes is used, this sheet should not be included in the request.			
Name and address: (Family name followed by given name; for a le designation. The address must include postal code and name of count address indicated in this Box is the applicant s State (that is, country) of residence is indicated below.)	try The country of the		
THORGAARD, Per Leonorevej 6 DK-9000 Aalborg Denmark	applicant and inventor inventor only (If this check-box		
	is marked, do not fill in below.)		
State (that is, country) of nationality: DK	State (that is, country) of residence:		
This person is applicant for the purposes of:	les of America only the Supplemental Box		
Name and address: (Family name followed by given name; for a le designation. The address must include postal code and name of coun- address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)	gal entity, full official try. The country of the of residence if no State This person is: applicant only		
KJÆRGAARD, Søren Christensen	applicant and inventor		
Nordvestvej 11 DK-9000 Aalborg			
Denmark	inventor only (If this check-box is marked, do not fill in below.)		
State (that is, country) of nationality:	State (that is, country) of residence: DK		
This person is applicant for the purposes of:	States except es of America the United States the States indicated in the Supplemental Box		
Name and address: (Family name followed by given name; for a le designation. The address must include postal code and name of coun address indicated in this Box is the applicant s State (that is, country) of residence is indicated below.)	gal entity, full official by. The country of the of residence if no State This person is: applicant only		
ANDREASSEN, Steen Kong Georgs Vej 7 DK-9000 Aalborg	applicant and inventor		
Denmark	inventor only (If this check-box is marked, do not fill in below.)		
State (that is, country) of nationality: DK	State (that is, country) of residence: DK		
This person is applicant for the purposes of: All designated the United States the United States and the United States	States except the United States the States indicated in the Supplemental Box		
Namc and address: (Family name followed by given name; for a le designation. The address must include postal code and name of coun address indicated in this Box is the applicant's State (that is, country) of residence is indicated below.)	try. The country of the		
	applicant and inventor		
	inventor only (If this check-box is marked, do not fill in below.)		
State (that is, country) of nationality:	State (that is, country) of residence:		
This person is applicant all designated all designated States except the United States of America only the States indicated in the States of America only the Supplemental Box			
Further applicants and/or (further) inventors are indicated on another continuation sheet.			

Box No.V DESIGNATION OF STATES					
The foll	lowing designations are hereby made under Rule 4.9(a) (m	ark	the app	olicable check-boxes; at least one must be marked):	
	al Patent				
X AP	AP ARIPO Patent: GH Ghana, GM Gambia, KE Kenya, LS Lesotho, MW Malawi, SD Sudan, SL Sierra Leone, SZ Swaziland, TZ United Republic of Tanzania, UG Uganda, ZW Zimbabwe, and any other State which is a Contracting State of the Harare Protocol and of the PCT				
X EA	EA Eurasian Patent: AM Armenia, AZ Azerbaijan, BY Belarus, KG Kyrgyzstan, KZ Kazakhstan, MD Republic of Moldova, RU Russian Federation, TJ Tajikistan, TM Turkmenistan, and any other State which is a Contracting State of the Eurasian Patent Convention and of the PCT				
	P European Patent: AT Austria, BE Belgium, CH and LI Switzerland and Licchtenstein, CY Cyprus, DE Germany, DK Denmark, ES Spain, FI Finland, FR France, GB United Kingdom, GR Greece, IE Ireland, IT Italy, LU Luxembourg, MC Monaco, NL Netherlands, PT Portugal, SE Sweden, and any other State which is a Contracting State of the European Patent Convention and of the PCT				
▼ OA	MOA OAPI Patent: BF Burkina Faso, BJ Benin, CF Central African Republic, CG Congo, CI Côte d Ivoire, CM Cameroon, GA Gabon, GN Guinea, GW Guinea-Bissau, ML Mali, MR Mauritania, NE Niger, SN Senegal, TD Chad, TG Togo, and any other State which is a member State of OAPI and a Contracting State of the PCT (if other kind of protection or treatment desired, specify on dotted line)				
Nation	al Patent (if other kind of protection or treatment desired, spec	ify o	m dotte	ed line):	
X AE	United Arab Emirates	X	LR	Liberia	
XAL	Albania	=		Lesotho	
X AM	Armenia	=		Lithuania	
I =	Austria and utility model			Luxembourg	
	Australia	_		Latvia	
I ===	Azerbaijan	=		Morocco	
1 —	Bosnia and Herzegovina	=		Republic of Moldova	
1 =	Barbados			Madagascar	
1 —	Bulgaria			The former Yugoslav Republic of Macedonia	
_	Brazil	فبتد		The former rugosiav Republic of Maccoonia	
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ĭ ES	Spain	=	SG	Singapore	
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Precautionary Designation Statement: In addition to the designations made above, the applicant also makes under Rule 4.9(b) all other designations which would be permitted under the PCT except any designation(s) indicated in the Supplemental Box as being excluded from the scope of this statement. The applicant declares that those additional designations are subject to confirmation and that any designation which is not confirmed before the expiration of 15 months from the priority date is to be regarded as withdrawn by the applicant at the expiration of that time limit. (Confirmation (including fees) must reach the receiving Office within the 15-month time limit.)					

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Sheet No.	

Box No. VI PRIORITY CLAIM Further priority claims are indicated in the Supplemental Box.						
Filing date Number Where earlier application is:			ion is:			
(day/month/year)		international application: receiving Office				
item (1) 03/02/1999	PA 19	99 00129 🗸	DK	V		
item (2)	1					
12/05/1999	PA 19	99 00649 🗸	DK	<i>V</i>		
item (3) 17/06/1999	PA 19	999 00859 🗸	DK	/		
The receiving Office is re of the earlier application(s) (only if	the earlier applic	ation was	filed with the	Office which for the), (2), and (3)
* Where the earlier application is	an ARIPO	application, it is ma	ndatory to	indicate in the Si	upplemental Box at least on	e country party to the Paris
Convention for the Protection of I Box No. VII INTERNATION		ARCHING AUT			ed (Rule 4.10(0)(11)). See St.	ірріетепій Вох.
Choice of International Searce (if two or more International Searce competent to carry out the intern	hing Autl	hority (ISA) Req	uest to u	se results of ear		to that search (if an earlier tional Searching Authority):
the Authority chosen; the two-letter			e (day/moni	h/year)	Number	Country (or regional Office)
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This international application		This internationa	al applicat	ion is accompa	nied by the item(s) mark	ed below:
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description (excluding	•	2. separate s	igned pov	ver of attorney		
sequence listing part) :	25	, – •	•	• •	reference number, if an	y:
claims :	11	4. statement	explainin	g lack of signat	ure	
abstract :	1	5. priority do	ocument(s	s) identified in F	Box No. VI as item(s):	
drawings :	10	_			tion into (language):	
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Jens Jørgen Schmidt For receiving Office use only						
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3. Corrected date of actual receipt due to later but timely received papers or drawings completing the purported international application:			received:			
4. Date of timely receipt of the required corrections under PCT Article 11(2):			not received:			
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PATENT COOPERATION TREATY

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REC'D	25	NUL	2001	
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INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference	FOR FURTHER ACTION	See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)
International application No.	International filing date (day mo	nth year) Priority date (day/month/year)
PCT/DK00/00040	01.02.2000	03.02.1999
International Patent Classification (IPC) o	r national classification and IPC7	
A 61 B 5/08, A 61 M 1	6/00	
Applicant REES, REESE, Stephan Edward	et al	
2. This REPORT consists of a total of This report is also accompanies a mended and are the best of the total	e applicant according to Article 36 of sheets, including the sheets of the sh	ng this cover sheet. the description, claims and/or drawings which have ontaining rectifications made before this Authority
These annexes consist of a total of		·
3. This report contains indications rel	ating to the following items:	
I Basis of the report		
II Priority	•	
III Non-establishment of	opinion with regard to novelty, in	ventive step and industrial applicability
. IV Lack of unity of inven	tion	
V Reasoned statement un citations and explanati	nder Article 35(2) with regard to a one supporting such statement	novelty, inventive step or industrial applicability;
VI Certain documents cite	ed	·
VII Certain defects in the i	nternational application	,
VIII Certain observations o	n the international application	,
Date of submission of the demand	Data of	completion of this report
	Date of	compositon of this report
07.08.2000	12.0	6.2001
Name and mailing address of the IPEA/SE	Authoriz	ed officer
Patent- och registreringsverket Box 5055 G-102 42 STOCKHOLH Facsimile No. 08-667 72 88 Form PCT/IPEA/409 (cover sheet) (January	Telepho	Bengtsson/EE ne No. 08-782 25 00

International application No.

PCT/DK00/00040

I. Bas	Basis of the report	
1. With	Vith regard to the elements of the international application:*	
	the international application as originally filed	
\boxtimes	the description:	
الحسا	nages 1 – 25	, as originally filed
	pages	, filed with the demand
	pages, filed wit	
\boxtimes	the claims:	
	pages	, as originally filed
	pages, as amend	
	pages	, filed with the demand
<u> </u>	pages <u>1-11</u> , filed with	h the letter of 27.04.2001
\boxtimes	the drawings:	
	pages 1-10	, as originally filed
	pages	, filed with the demand
	pages, filed with	the letter of
لــا	the sequence listing part of the description:	
	pages	, as originally filed
	P-6-0	, filed with the demand
2 With r	, , , , , ,	
uic mi	ith regard to the language, all the elements marked above were available or furner international application was filed, unless otherwise indicated under this item.	
These	ese elements were available or furnished to this Authority in the following lang	uage which is:
	the language of a translation furnished for the purposes of international sear	ch (under Rule 23.1(b)).
	the language of publication of the international application (under Rule 48.3	
	the language of the translation furnished for the purposes of international pr	
	01 33.3).	
3. With representation	th regard to any nucleotide and/or amino acid sequence disclosed in the internal int	national application, the international
	contained in the international application in written form.	
=	filed together with the international application in computer readable form.	
	furnished subsequently to this Authority in written form.	
==	furnished subsequently to this Authority in computer readable form.	
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	International application as filed has been furnished	
	The statement that the information recorded in computer readable form is id been furnished.	entical to the written sequence listing has
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ا ا	The amendments have resulted in the cancellation of:	
	the description, pages	
. [the claims, Nos.	
[the drawings, sheet/fig	:
5. []	This report has been established as if (some of) the amendments had not bee beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 7)	n made, since they have been considered to go 0.2 (c)).**
111 11113	placement sheets which have been furnished to the receiving Office in response this report as "originally filed" and are annexed to this report since they do not I 70.17).	to an invitation under Article 14 are referred to contain amendments (Rules 70.16
* .Any rej	y replacement sheet containing such amendments must be referred to under iten	ı l and annexed to this report.
	T/DE A/400 (Boy I) (Improved 1000)	·

International application No.

PCT/DK00/00040

V.	Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial ap	nlicability
	citations and explanations supporting such statement	pricability,

1. Statement

Novelty (N) Claims 1-50 YES Claims NO Inventive step (IS) Claims YES Claims NO. Industrial applicability (IA) Claims YES Claims NO

2. Citations and explanations (Rule 70.7)

The claimed invention relates to an automatic lung parameter estimator, which has a gas-mixing unit supplying a first homogeneous gas to the inlet of a ventilator via a gas mixer. A second gas is mixed with the first gas in the mixer. A computer determines respiratory parameters and controls the gas mixture supplied to the patient.

The following relevant documents were cited in the International Search Report:

D1: EP 0753320, A1 D2: US 5103814, A D3: EP 0502270, A1

Document D1, which is the closest prior art, describes artificial ventilation system. The system comprises respiratory gas delivery unit with а regulatory unit connectable to the lung of the patient and a parametermonitoring unit comprising a blood gas analyser.

The documents D2 and D3 are two types of respirator system, which both differ significantly from the invention.

The cited document D2 relates to a self-compensating patient respirator. The ventilator comprises a respirator and pulse-oximeter for determining the body oxygen saturation of a patient. An oxygen comparison circuit compares the body oxygen saturation to a predetermined body oxygen saturation level. A control is coupled to the respirator, for periodically decreasing the percentage of oxygen in a respiratory gas within the ventilator while the body oxygen saturation is greater than the predetermined body oxygen saturation level.

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International application No.

PCT/DK00/00040

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Continuation of: V

Document D3 relates to tissue oxygen measuring system, which has a ventilation unit, a control unit for controlling the gaseous content of the air and an arrangement for producing trigger signals. A measurement system is included for measuring oxygen in the tissue and supplying data regarding the measured results, and a data processing unit computing information regarding blood flowing in the tissue.

D1 a device is known for determining respiratory parameters. The parameters that are determined in D1 are either settings for the artificial ventilation system or functions of the measured parameters over time. No parameters in D1 being descriptive of the pulmonary gas exchange are computed as the purpose of the artificial ventilation system. Disclosed is to obtain an optimised artificial ventilation of a lung system while preventing injuries of the lungs. The invention according to claims 1-50 is therefore new and involves an inventive step.

The invention according to claims 1--50 is novel (N) and is considered to involve an inventive step (IS). The invention according to claims 1--50 is considered to have industrial applicability (IA).

PCT/DK00/00040

AMENDED SET OF CLAIMS

REPLY TO FIRST WRITTEN OPINION 27 APRIL 2001

5 1. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction
different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2, FEO2, PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

- the computer being adapted for retrieving and storing at least two measurements being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the at least two measurements being conducted at respective levels of oxygen in the gas flow passing into the respiratory system, the computer further being adapted for determining at least two respiratory
- parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the pulmonary gas exchange of the individual, the determination being based on the at least two measurements.

- 2. A device according to claim 1, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 5 3. A device according to claim 1 or 2, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising

determining, based on at least two measurements, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection 10 means,

producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

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4. A device according to any of claims 1-3, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the

25 respiratory system,

the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously.

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5. A device according to claim 4, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.

- 6. A. device according to any of claims 1-5, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item
 5 accordingly if said parameter exceeds said threshold value.
- 7. A device according to any of claim 1-6, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2,
 SpO2, PaO2, PpO2) and produce a control data item accordingly.
- 8. A device according to claim 7, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurements.
 - 9. A device according to claim 7, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 10. A device according to any of claims 7-9, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 11. A device according to any of claims 1-10, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.
 - 12. A device according to any of claims 1-11, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 13. A device according to any of the preceding claims, wherein the oxygen saturation in35 the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.

14. A device according to any of claims 1-13, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

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15. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2)

20 in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2,

PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer being further adapted for performing a procedure at least once, the procedure comprising

determining, based on data stored within the data structure, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection means,

producing a possible control data item based on the target, and

retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

5 16. A device according to claim 15, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure in data storage means associated with the computer, in which the stored outputs are mutually related and related to the output from the first detection means and the second detection means, and the output from the four detection means can be retrieved simultaneously.
 - 17. A device according to claim 16, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 18. A device according to claim 15 or 16, wherein the computer is adapted for determining at least one respiratory parameter (Rdiff, shunt, V/Q, H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 30 19. A device according to claim 18, wherein at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.

- 20. A device according to claim 18 or 19, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 5 21. A. device according to any of claims 15-20, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.

22. A device according to any of claims 15-21, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

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23. A device according to claim 22, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).

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- 24. A device according to claim 22, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 25 25. A device according to any of claims 22-24, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.

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26. A device according to any of claims 15-25, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

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- 27. A device according to any of claims 15-26, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 28. A device according to any of claims 15-28, wherein the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
 - 29. A device according to any of claims 15-28, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

30. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2)

in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2, FEO2,

PIO2, PE'O2, FĒO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing at least a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer further being adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from

the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

- 31. A device according to claim 30, wherein the assessment of change in oxygen level in
 5 the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).
- 32. A device according to claim 30, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 33. A device according to any of claims 30-32, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one
 gas, in response to said control data item from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 34. A device according to any of claims 30-33, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising
- determining, based on at least one measurement, whether additional measurements are required,
 - asserting a possible desired target defining a desired output of the first detection means,
- producing a possible control data item based on the target, and
 retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.
- 35. A device according to any of claims 30-34, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- 5 the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously.
- 36. A device according to claim 35, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 37. A device according to any of claims 30-36, wherein the computer is adapted for
 15 determining at least one respiratory parameter (Rdiff, shunt, V/Q, H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 38. A device according to claim 37, wherein at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.
 - 39. A device according to claim 37 or 38, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 40. A. device according to any of claims 30-39, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.
 - 41. A device according to any of claims 30-40, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

- 42. A device according to any of claims 30-41, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 5 43. A device according to any of claims 30-42, wherein the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
- 44. A device according to any of claims 30-42, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.
 - 45. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is an apparently healthy individual.
 - 46. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is considered to have a risk of suffering from hypoxemia.
- 47. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is suffering from hypoxemia.
- 48. Method according to claim 47, wherein the individual is suffering from one or more disease(s) selected from the group(s) comprising left sided heart failure, adult respiratory distress syndrome, pneumonia, postoperative hypoxemia, pulmonary fibrosis, toxic pulmonary lymphoedema, pulmonary embolisms, chronic obstructive pulmonary disease and cardiac shunting.
- 49. A computer system comprising at least one general purpose computer having one or more computer programs stored within data storage means associated therewith, the computer system being arranged for as well as being adapted for determining one or more respiratory parameters according to any of claims 1-48.

50. A computer program product being adapted to enable a computer system comprising at least one general purpose computer having data storage means associated therewith and being arranged suitably to determine one or more respiratory parameters according to any of claims 1-48.

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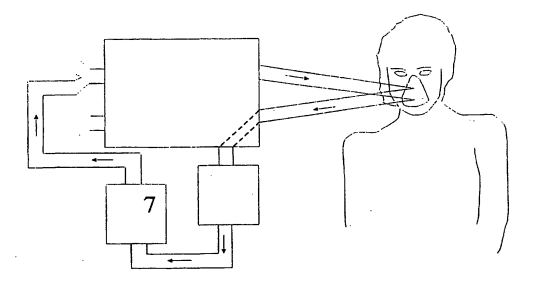
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(57) Abstract

A device for determining one or more respiratory parameters relating to an individual is disclosed, as well as a method for determining one or more respiratory parameters by means of the device, wherein the individual is suffering from hypoxemia or is at risk of hypoxemia. However, the method and the device may also be applied to healthy individual e.g. for testing of medicaments. The device is controlled by a computer equipped with suitable software and includes functionality for on-line continuous data collection, automatic assessment of the timing of measurements, automatic assessment of the next target (oxygen saturation of arterial blood (SpO2)), automatic assessment of the appropriate fraction of oxygen in inspired gas (FIO2) settings to achieve the target SpO2, automatic control of the FIO2, on-line parameter estimation, and automatic assessment of the number of measurements required.

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AUTOMATIC LUNG PARAMETER ESTIMATOR

The present invention relates to a device for determining one or more respiratory parameters relating to an individual. The device may include functionality for on-line continuous data collection, automatic assessment of the timing of measurements, automatic assessment of the next target (oxygen saturation of arterial blood (SpO2)), automatic assessment of the appropriate fraction of oxygen in inspired gas (FlO2) settings to achieve the target SpO2, automatic control of the FlO2, on-line parameter estimation, and automatic assessment of the number of measurements required. This functionality is achieved through a novel device including ventilatory equipment, blood gas analysis equipment and computer hardware and software.

Furthermore, the present invention relates to a method for determining one or more respiratory parameters by means of the above-mentioned device, wherein the individual is suffering from hypoxemia or is at risk of hypoxemia. The individual may also be a healthy individual.

The use of the device for examination and monitoring respiratory parameters relating to humans are of particular interest, but the device may also be applied to farm animals such as pigs, or to domestic animals such as dogs.

BACKGROUND

Oxygen enters the body with inspiration and diffuses from the lungs into the blood.

Subsequently the blood circulation transports oxygen to the tissues. Disorders of oxygen transport from the inspired air into the blood can result in a low oxygen saturation of the blood. These disorders in oxygen uptake include abnormal ventilation of the lung, seen in for example chronic obstructive pulmonary disease; abnormal oxygen diffusion in the lung, seen in for example pulmonary fibrosis; and abnormal perfusion (i.e. blood flow) through the lung. Estimation of parameters describing these oxygenation problems is important for diagnosis, monitoring and assessing appropriate therapeutic intervention. This is true in a wide variety of patients, from those who are automatically ventilated and who often require continuous supplement of oxygen, to out-patients who only suffer from dyspnoe during exercise.



In clinical practice the clinician usually relies upon simple measurements or variable estimates to assess the patients oxygenation problems. These include qualitative estimates obtained from stethoscopy or chest X-ray. They also include more quantitative estimates such as arterial oxygen saturation, the alveolar-arterial oxygen pressure gradient, or estimates of the "effective shunt", a parameter which describes all oxygenation problems in terms of a fraction of blood which does not flow through the lungs (Siggaard-Andersen and Siggaard-Andersen, 1985).

Whilst the "effective shunt" is a parameter which has been used widely in the clinical literature it cannot adequately describe the 'clinical' picture seen in patients when the inspired oxygen fraction is varied. This observation is illustrated in Figure 1 where the "effective shunt" has been estimated for a single patient at four different inspired oxygen fractions, and varies from 15-25%.

In contrast to the poor clinical description of oxygenation problems, detailed experimental techniques such as the Multiple Inert Gas Elimination Technique (MIGET) (Wagner et al., 1974) have been developed which describe the parameters of models with as many as fifty lung compartments. The parameters of these models give an accurate physiological picture of the patient. Whilst the MIGET has found widespread application as an experimental tool its use as a routine clinical tool has been somewhat limited (Wagner et al., 1987). This is largely due to the cost and complexity of the technique.

As stated previously, "effective shunt" is insufficient to describe oxygenation problems. Further parameters describing the patient's oxygenation problem can be obtained from data where inspired oxygen is varied, i.e. data similar to that presented in Figure 1. This was first recognised by Riley et al. (1951a, 1951b) and later by King et al. (1974). These authors used mathematical models to divide the oxygenation problem into that due to an alveolar-lung capillary drop in the partial pressure of oxygen, and that due to a shunt problem. To estimate two parameters describing the oxygenation problem requires taking measurements of blood samples and of ventilatory variables at each inspired oxygen fraction. Estimating lung parameters using the data from four inspired oxygen fractions required four blood samples, a procedure which is still rather time consuming and in some environments impractical.

More recently, development of non-invasive methods for measuring the oxygen saturation of the blood have lead to renewed interest in estimation of parameters describing oxygen transport obtained by varying FIO2. Andreassen et al. (1996, 1999), Sapsford et al. (1995), de Gray et al. (1997) and Roe et al. (1997), have presented the use of two parameter mathematical models of oxygen transport, the oxygenation problem being described as shunt combined with either a diffusion abnormality (Andreassen et al. (1996, 1999)) or due to a ventilation/perfusion (\vec{V}/\vec{Q}) mismatch (Sapsford et al. (1995), de Gray et al (1997), Roe et al., (1997)). These model representations have been shown to provide identical fits to routine blood gas and ventilatory data obtained by varying FIO2 (Rees et al. 1997).

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The clinical relevance of the two parameter models is illustrated in Fig.2, where increases in the pulmonary shunt parameter results in a vertical depression of the FIO2/ SaO2 curve, (V-shift) and abnormalities in the second parameter (ventilation/perfusion (\dot{V}/\dot{Q}) mismatch or oxygen diffusion resistance (Rdiff)) results in a lateral displacement of the FIO2/ SaO2 curve. Clearly, the lateral displacement of the FIO2/ SaO2 curve (H-shift) is clinically a more significant problem as it describes a situation where large changes in oxygen saturation can occur for only small changes in FIO2. In this situation the patient is at increased risk of an oxygenation problem.

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The two parameter model of Sapsford et al. (1995), has been shown to fit data from normal subjects; patients before and after thoracotomy (Sapsford et al. 1995, de Gray et al., 1997); and patients during (Sapsford et al. 1995, Roe et al., 1997), and after (Roe et al., 1997) abdominal surgery. Similarly, the two-parameter model described by 25 Andreassen et. al. has been shown to fit data from normal subject and postoperative cardiac patients (Andreassen, 1999) and a wide range of as yet un-published results. Examples of these results are shown in Fig. 3.

In contrary to detailed experimental approaches (e.g. the MIGET), these two parameter models can be used routinely in clinical practice. In particular, these techniques may find application in the monitoring and choice of therapeutic treatment for patients with left-sided heart failure, or to assess patients risk of post-operative hypoxaemia.

Until now, estimation of oxygenation parameters has involved manual titration of the FIO2/ 35 SaO2 curve and off-line estimation of the parameter values. This is time consuming with experimental times of approximately 45 minutes, not including the time required for off line parameter estimation. This limits the use of the method as a clinical tool.

DESCRIPTION OF THE INVENTION

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It is an object of the present invention to provide a device for estimation of one or more respiratory parameters including oxygenation parameters and lung parameters relating to an individual in which the necessary quantities for enabling an estimation of respiratory parameters are collected automatically by a computer of the device so as to provide an automated estimation of said parameters.

It is a further object to provide a device wherein the necessary measurements at varying oxygen levels are obtained in an at least semi-automated manner whereby the experimental time for said estimation may be reduced. By reducing the procedural time these techniques have potential for routine clinical use.

It is a still further object to provide a device which is adapted for assessing a possible new target of the level of oxygen in the blood circulation based on the previously obtained measurement(s).

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It is a yet still further object to provide a device, which is adapted for assessing an appropriate change in the current level of oxygen in the inspired gas to obtain a given target of the level of oxygen in the blood circulation.

25 The use of the device on humans is of particular interest, but the device may also be applied to farm animals such as pigs, or to domestic animals such as dogs.

The device might be of value in all kind of patients in which hypoxemia occurs or may occur. These conditions may e.g. be selected from the group comprising left sided heart failure, adult respiratory distress syndrome, pneumonia, postoperative hypoxemia, pulmonary fibrosis, toxic pulmonary lymphoedema, pulmonary embolisms, chronic obstructive pulmonary disease and cardiac shunting.

Thus, the present invention relates in a first aspect of the present invention to a device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas.

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FE'O2, FĒO2, PĒO2, PĒO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly, the computer being adapted for retrieving and storing at least two measurements being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the at least two measurements being conducted at respective levels of oxygen in the gas flow passing into the respiratory system, the computer further being adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on the at least two measurements.

Hence, in its broadest aspect, the invention relates to a device for determining one or more respiratory parameters relating to an individual. By the term "individual" is herein understood an individual selected from the group comprising humans as well as farm animals, domestic animals, pet animals and animals used for experiments such as monkeys, rats, rabbits, etc.

By the term "respiratory parameters" is herein understood parameters relating to oxygen transport from the lungs to the blood, such as parameters related to abnormal ventilation,

resistance to oxygen uptake from the lungs to the lung capillary blood, and parameters related to shunting of venous blood to the arterial blood stream. These respiratory parameters may be given as absolute values or relative values as compared to a set of standard values and the parameters may further be normalised or generalised to obtain 5 parameters that are comparable to similar parameters measured for other individuals, at least for individuals of the same species.

Thus, the computer may further be adapted for determining at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the 10 individual, and said parameter(s) (Rdiff, shunt, V/Q, H-shift, V-shift) may alternatively or additionally be generalised parameters being comparable to similar parameter(s) determined for other individuals.

In a preferred embodiment, the computer of the device is further adapted for performing a 15 procedure at least once, the procedure comprising

determining, based on at least two measurements, whether additional measurements are required.

asserting a possible desired target defining a desired output of the first detection means,

producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means. The control data item produced thereby may be outputted to a human operator by means of an output device so that the operator can adjust the level of oxygen in the 25 inspired gas flow. Alternatively, the control data item may be used by another part of or a computer program within the computer or by an external control device for automatically control of the means for controlling the flow to the gas-mixing unit of at least one gas.

According to a preferred embodiment of the present invention, the second detection 30 means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FEO2, PE'O2, PEO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

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5 the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously. This/these measurement(s) enable(s) the computer to estimate or establish the oxygen consumption of the individual, either
10 implicitly as part of the estimation of respiratory parameters, or the computer may further be adapted for explicitly establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.

It is advantageous for the device according to the present invention that the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value. By determining whether an equilibrium state of the individual is obtained the timing of the steps of the procedure can be controlled efficiently and the overall time for performing the procedure may be further reduced.

It is also advantageous if the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly so that the oxygen level can be adjusted according to the data item. The actual adjustment may be performed by an operator of the device, in which case the data item is outputted to an output device. Alternatively and preferably the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly. The data item may instead be outputted to an external device, which is suitable for performing an automated control of the control means so as to adjust the oxygen level accordingly.

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The assessment of change in oxygen level in the inspired gas may in an embodiment of the invention be based on a predefined set of data representing statistical distributions of variables stored within data storage means associated with the computer and on said measurements. Details of how this may be performed are disclosed in the detailed description of the invention. Alternatively, the assessment of change in oxygen level in the inspired gas may be based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow. Typically, the oxygen level is changed stepwise or following a ramp function and the change over time of the oxygen level in the blood circulation or the level of oxygen in the expired gas is monitored. However, monitoring of another gas, such as CO₂, or another variable of the patient may additionally or alternatively be employed.

It is preferred that one gas is atmospheric air and that another of the gasses is more or less pure oxygen, i.e. has an oxygen fraction higher than that of atmospheric air,

- preferably in the range 0.85 to 1.00. Alternatively or additionally, another gas may be supplied which has an oxygen fraction below that of atmospheric air, i.e. in the range of 0.00 to 0.21, preferably of 0.00 to 0.05. Thereby the oxygen level of the inspired gas may be varied not only to level above that of atmospheric air but also below that level, thus providing a wide range of possible levels for performing measurements of the individual.
- The gas having a low oxygen fraction may be supplied from a source of more or less pure nitrogen N₂ or another suitable physiologically neutral gas, such as helium H₂, or it may be re-circulated expired gas from the individual, preferably after reduction of the level of CO₂ in the expired gas.
- The device should ensure by means of a security arrangement that the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably for human beings in the range of 85 to 100% to avoid the risk of damage to organs. This condition varies for different species of animals.
- 30 The first detection means is preferably arranged for detecting a variable relating to the saturation level of oxygen in the arterial blood stream by means of an invasive or a non-invasive technique, which latter is preferred. Thus, the first detection means is in an advantageous embodiment a pulse oximeter. Alternatively, the level of oxygen in the venous blood stream may be measured by means of an invasive or a non-invasive technique, the latter again being the preferred one.

According to a second aspect, the present invention relates to a device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening.

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and 10 having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2,

PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer being further adapted for performing a procedure at least once, the procedure comprising

determining, based on data stored within the data structure, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection means,

producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

accordingly, and

10

According to a third aspect, the present invention relates to a device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2)

15 in the blood circulation of the individual and producing an output to the computer

second detection means for detecting the level of oxygen (FIO2, FEO2,

PIO2, PE'O2, FEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing at least a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer further being adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

The second aspect as well as the third aspect of the invention is disclosed above in the most fundamental embodiment which according to the present invention may be combined with the additional features disclosed above with relation to the first aspect of the invention.

The device may be used to obtain and/or compare one or more respiratory parameters relating to one or more individual(s). The individual may be a healthy individual, at risk of suffering from hypoxemia, or suffering from hypoxemia.

By the term "the individual is at risk of suffering from hypoxemia" is herein understood that the individual has a higher/increased risk of suffering from hypoxemia compared to a healthy individual. The increased risk of suffering from hypoxemia may e.g. be due to a hereditary predisposition, a post-operative condition and/or various diseases.

By the term "hypoxemia" is herein meant that the oxygen saturation in the blood from the individual is below 92%. Examples of diseases that can cause hypoxemia are left sided heart failure, adult respiratory distress syndrome, pneumonia, postoperative hypoxemia, pulmonary fibrosis, toxic pulmonary lymphoedema, pulmonary embolisms, chronic obstructive pulmonary disease and cardiac shunting.

The present invention also relates to a computer system comprising at least one general purpose computer having one or more computer programs stored within data storage means associated therewith, the computer system being arranged for as well as being adapted for determining one or more respiratory parameters according to the devices and/or methods disclosed above.

Furthermore, the present invention relates to a computer program product being adapted to enable a computer system comprising at least one general purpose computer having data storage means associated therewith and being arranged suitably to determine one or more respiratory parameters according to the devices and/or methods disclosed above.

GLOSSARY

2	5	
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	FIO2	Fraction of oxygen in inspired gas.
	PIO2	Pressure of oxygen in inspired gas.
	SaO2	Oxygen saturation of arterial blood, measured from a blood sample.
	PaO2	Pressure of oxygen in arterial blood, measured from a blood sample.
30	SpO2	Oxygen saturation of arterial blood, measured transcutaneously.
	PpO ₂	Pressure of oxygen in arterial blood, measured transcutaneously.
	FĒCO2	Fraction of carbon dioxide in the mixed expired gas.
	FE'O2	Fraction of oxygen in expired gas at the end of expiration.
	FĒO2	Fraction of oxygen in the mixed expired gas.

	PĒCO2	Pressure of oxygen in the mixed expired gas.
	PE'O2	Pressure of oxygen in expired gas at the end of expiration.
	Vt	Tidal volume, i.e. volume of gas breathed per breath.
	f	Respiratory frequency, i.e. number of breaths per minute.
5	VO2	Oxygen consumption, i.e. the amount of oxygen consumed by the tissues per minute.
	Vd	Dead space i.e. the volume of the lung not involved in exchanging gases with the blood.
10	shunt	Respiratory parameter representing the faction of blood not involved in gas exchange.
,0	Rdiff	Respiratory parameter representing a resistance to oxygen diffusion across the alveolar lung capillary membrane.
	\dot{V}	Ventilation.
	\dot{V} / \dot{Q}	Respiratory parameter representing the balance between ventilation and
15		perfusion in a region of the lung.
	V-shift	Respiratory parameter representing a vertical shift in plots of FIO2
		against SaO2 , FIO2 against SpO2, FE'O2 against SaO2, or FE'O2
		against SpO2 .
	H-shift	Respiratory parameter representing a horizontal shift in plots of FIO2
20		against SaO ₂ , FIO ₂ against SpO ₂ , FE'O ₂ against SaO ₂ , or FE'O ₂ against SpO ₂ .

BRIEF DESCRIPTION OF THE FIGURES

25 Fig.1. Plot of the inspired oxygen fraction (FIO2, x-axis) against the arterial oxygen saturation (SaO2, SpO2, y-axis) for 1 patient. For each data point (A-D) the "effective shunt" has been estimated from a single parameter shunt model (Siggard-Andersen and Siggaard-Andersen 1985), giving values of point A = 15%, point B = 15%, point C = 20%, point D = 25%.

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Fig. 2. Plots of the inspired oxygen fraction (FIO2, x-axis) against model predicted arterial oxygen saturation (SaO2, SpO2, y-axis) for 1) a normal subject with shunt = 5% and Rdiff = 0 kPa/(I/min) (solid line), 2) a hypothetical patient with a Rdiff or ventilation/perfusion disorder (dotted line), and 3) a hypothetical patient with a shunt disorder (dashed line).

Line A illustrates the vertical displacement of the curve (V-shift) due to a shunt disorder, whilst line B illustrates the horizontal displacement of the curve (H-shift) due to a ventilation perfusion of oxygen diffusion abnormality.

- Fig. 3. Plots of the inspired oxygen fraction (FIO2, x-axis) against arterial oxygen saturation (SaO2, SpO2, y-axis). Each of the vignettes illustrates data (crosses) and model predicted curves fitted, to this data from: A a normal subject (shunt = 5%, Rdiff = -1.5 kPa/(I/min)), B a post-operative cardiac patient (shunt = 9.5%, Rdiff = 81.0 kPa/(I/min)), C a post-operative hysterectomy patient (shunt = 7%, Rdiff = 15.2
 kPa/(I/min)), D a poorly compensated cardiac patient (shunt = 15%, Rdiff = 22.9 kPa/(I/min)), and E a patient residing in the intensive care unit (shunt = 7%, Rdiff = 31.0 kPa/(I/min)).
- Fig. 4. Experimental set-up working with nitrogen for subathmospheric oxygen levels. The system includes: 1) A Gas Delivery Unit including gas inlets (1a, 1b), a gas mixer (1c), a flow or pressure gradient (1d), and equipment for the measurement and/or setting of inspired oxygen fraction (FIO2), tidal volume and respiratory frequency (1e); 2) Equipment for measurement of expired gases including an oxygen monitor placed so as to measure end tidal oxygen fraction (2a), and/or an expiratory reservoir, used with an oxygen monitor and/or a carbon dioxide monitor to measure the fraction of gas in or leaving the expiratory reservoir (FĒO2, FĒCO2) (2b); 3) Measurement of arterial oxygen saturation (SaO2) via e.g. a pulse oxymeter (SpO2); 4) Measurements of arterial or venous blood gas samples (optional); 5) Measurement of cardiac output (optional); 6) A computer system including software for automatic collection of data (6a), monitoring the steady state of the patients/subjects oxygenation (6b), a feedback controller for adjusting inspired oxygen fraction (6c), and estimation of gas exchange parameters. Dashed arrowed lines illustrate the flow of information to the computer. Dotted arrowed lines illustrated the control of the gas delivery unit by the computer.
- 30 Fig. 5. Experimental set up using a rebreathing technique for subatmospheric oxygen levels. Figure 5 illustrates a modification to the set-up of Figure 4. It includes all other components illustrated in Figure 4, plus a carbon dioxide removal device to eliminate carbon dioxide from the re-inspired gases (box 7). All other points 1-6 are the same as Fig. 4.



- A: Begin parameter estimation if FIO2>1.00 and SpO2>0.85
- B: Continuous data recording from gas delivery unit, pulse oxymeter and expiratory gas measurement devices.
- 5 C: Set oxygen level (FIO2).
 - D: Monitor O2 equilibrium.
 - E: Equilibrium level.
 - F: Record measurement.
 - G: Sufficient number of measurements?
- 10 H: Estimate new FIO2.
 - I: Estimate Pulmonary Parameters.
 - Fig.7. (algorithm 1) Assessing whether another measurement is necessary and determining the target SpO2 for that measurement. If current FIO2 = 1.00 and SpO2 <
- 15 0.85% do not perform measurement.
 - A: Is there 1 measurement of (SpO2) 1 where $0.85 \le (SpO2) 1 < 0.92$?
 - B: Target SpO2: $0.85 \le (SpO2) \ 1 < 0.92$
 - C: Was FIO2 = 1.00 at this measurement?
 - D: Patient too sick for measurement.
- 20 E: Is there 1 measurement of (SpO2) 2 where 0.92 ≤ (SpO2) 2 < 0.95?
 - F: Target SpO2: $0.92 \le (SpO2) \ 2 < 0.95$
 - G: FIO₂ = 1.00 at this measurement?
 - H: Target SpO2: (SpO2) $1 \le SpO2 < (SpO2) 2$
 - I: Is there 1 measurement of (SpO2) 3 where $0.95 \le (SpO2) 3 < 0.98$?
- 25 J: Target SpO2: 0.95 ≤ (SpO2) 3 < 0.98
 - K: Was FIO2 = 1.00 at this measurement?
 - L: Target SpO2: (SpO2) $2 \le$ SpO2< (SpO2) 3
 - M: Set $FIO_2 = 1.00$.
- 30 Fig. 8 (algorithm 2) This controller uses a mathematical model of oxygen transport with two parameters, shunt and either diffusion resistance or \dot{V}/\dot{Q} mismatch. Parameters are implemented as stochastic variables and as such have a probabilistic distribution.
 - A: Select appropriate a priori estimates for parameters

The patients lung parameters are represented as stochastic variables with probability distributions. These parameters need to be initialised with *a priori* distributions. If the patients lung parameters have been investigated previously, or if the patient belongs to a well-defined population there may be well-defined *a priori* distributions for the patient's lung parameters.

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- B: Target SpO2 = first target level
- C: Update parameter estimates with measurement data.
- 10 This is a Bayesian update of the parameter estimates for the measured values. The output of this process being revised probability distributions for the patients' lung parameters.
 - D: Is the parameter probability mass distributed within range.
- 15 If the probability distributions for the patients' lung parameters have a very narrow distribution, then they are estimated with good precision, and no further FIO2 settings or measurements are required.
- E: Predict SpO2 (distribution) when FIO2 lowered/raised by a predetermined percentage, using parameter estimates. The predetermined percentage is dependent on the conditions and the patient. The mathematical models can be used to predict the effects of varying FIO2 giving the current estimate of the probability distributions for the patients' lung parameters. Predictions can be obtained in terms of the probability of a certain oxygen saturation of the blood.

25

F: Is 10 % of probability mass < target SpO2.

If the predicted probability distribution for SpO2 is distributed evenly about the target SpO2 then the FIO2 is selected for the next measurement.

- 30 G: Set the selected FIO2 level.
 - H: Continue the algorithm only if there are more target SpO2 levels?
 - I: Set the next target SpO2 level.

Fig. 9 illustrates a graph of a patients parameter (A, x-axis) plotted against the probability that this parameter takes a certain value (P(A), y-axis). One of these graphs is used for each patient parameter (i.e. shunt, Rdiff and or V/Q). Before a measurement procedure begins an *a priori* distribution is obtained for each of the patient parameters from computer storage. Subsequently, these *a priori* estimates are updated as measured data presents. Typical distributions of the shunt parameter are illustrated for a normal healthy subject both a priori (solid line, mean shunt = 5%), and following update of the distribution with measured data (dashed line).

Fig. 10 illustrates model predicted arterial oxygen saturation (SaO2, SpO2, y-axis) when varying inspired oxygen fraction (FIO2, x-axis). Points A and B are measured FIO2/SpO2 values which are used to update parameter values (i.e. P(parameters | measurements)). The updated parameter values are then used to predict the change in SpO2 on varying FIO2 (i.e. P(SpO2 | FIO2)). These predictions are illustrated for two different FIO2 levels (C and D) and are plotted as probability distributions. The appropriate FIO2 level is then selected so that ≤ x% (in this case 10%) of the probability distribution is below the target SpO2 level (E).

DETAILED DESCRIPTION OF THE INVENTION

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The following description of preferred embodiments of the invention will focus on a device for automating the estimation of lung parameters. This device (Automatic Lung Parameter Estimator = ALPE) enables reduction in the time taken to obtain estimates of oxygenation parameters, with the total time including on-line estimation of parameters taking 10-15 minutes. By reducing the procedural time these techniques have potential for routine clinical use. This is only possible because of the substantial novelty in the ALPE which may include functionality for:

- 1) On-line continuous data collection
- 30 2) Automatic assessment of the timing of measurements
 - 3) Automatic assessment of the next target SpO2
 - Automatic assessment of the appropriate FIO2 settings to achieve the target SpO2
 - 5) Automatic control of the FIO2
 - 6) On-line parameter estimation
- 35 7) Automatic assessment of the number of measurements required

This functionality is achieved through a novel apparatus including ventilatory equipment, blood gas analysis equipment and computer hardware and software as described below.

5 Description of the Automatic Lung Parameter Estimator (ALPE): The Automatic Lung Parameter Estimator (ALPE) illustrated in Figure 4 may be used to assess oxygenation parameters in any patient, with these parameters being useful for diagnostic or monitoring purposes. Monitoring of patients' lung parameters is of particular value for those patients with ongoing treatment for example those patients artificially 10 ventilated or those receiving therapies for left-sided heart failure.

The ALPE can automatically determine the parameters of models of oxygen transport.

These parameters are obtained from numerous measurements including the FIO2/SpO2 curve, with this curve being constructed automatically by the apparatus for SpO2 varying between 0.85 to 1.00.

ALPE illustrated in Fig. 4 includes the following (numbers before paragraphs refer to the numbers in Figure 4):

- A Gas Delivery Unit This equipment includes: Two or more gas inlets, shown here delivering a) oxygen or nitrogen, and b) air; c) A gas mixer capable of mixing two input gases to the required fraction or concentration; d) A means of delivering the gases to the patient/subject i.e. a flow or pressure gradient; e) Equipment for the measurement and/or setting of inspired oxygen fraction (FIO2), tidal volume and respiratory frequency (or minute volume). The gas delivery unit included in the system can either be a stand-alone device offering only this functionality, or any other device, which includes this functionality such as patient ventilation devices (respirators) commonly used for intensive care patients. Ventilatory gases are delivered to and removed from the patient/subject through a face mask, mouth piece combined with a nose clip, laryngeal endotracheal tube etc.
 - 2) Measurement of expired gases Expired gases are measured using either: a) An oxygen monitor, placed so as to measure expiratory gases and sensitive enough to give measurement of the end tidal oxygen fraction (FE'O2), i.e. the fraction of oxygen in the expired gases at the end of an expiration. b) An expiratory reservoir.

- 3) Measurement of arterial oxygen saturation (SaO2) via e.g. a pulse oxymeter (SpO2).
- Measurements of arterial or venous blood gas samples may be taken or may be
 monitored continuously by invasive means and put manually into the system.
 These measurements are optional.
 - 5) Measurement of cardiac output may be put manually into the system. This measurement is optional.

15

- 6) A computer system including software for
 - a) Automatic collection of data from the gas delivery unit (FIO2, Vt, f), the expired gas measurement devices (FE'O2, FĒO2, FĒCO2 (optional)), and the pulse oxymeter (or any other measure of SpO2 or SaO2).

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- b) Monitoring the steady state of the patients/subjects oxygenation.
- c) A feedback controller, which determines whether a further measurement is required and automatically adjusts the inspired oxygen fraction to the most appropriate level.
 - d) Estimation of gas exchange parameters from the data collected.

Dashed arrowed lines on Figure 4 illustrate the flow of information to the computer. Dotted arrowed lines illustrated the control of the gas delivery unit by the computer.

A modification to the system is also included as part of this patent (Fig. 5). For environments where nitrogen (N2) or another physiologically neutral gas is not available the oxygen content of inspired gases can be reduced lower than air (FlO2air = 21%) by

re-breathing expired gases. In this situation, in addition to all other components illustrated in Figure 4 a carbon dioxide removal device is included in the system to eliminate carbon dioxide from the re-inspired gases (box 7 Figure 5). All other points 1-6 described above are the same as Figure 4.

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DETAILED DESCRIPTION OF THE FLOWCHARTS

The flowcharts are provided solely to illustrate the invention by reference to specific embodiments. These flowcharts and the algorithms included herein, while illustrating certain aspects of the invention, do not portray the limitations or circumscribe the scope of the disclosed invention.

Fig. 6 is a flowchart illustrating the processes involved during operation of the ALPE.

15 Box A: After set-up of the equipment as illustrated in Fig. 4 and 5 the parameter estimation procedure begins.

Box B: As part of this process the computer continuously collects data from the other equipment, including FIO2 and SpO2 (and/or FE'O2, Vt, f, FĒO2, FĒCO2).

20

Box C: An initial inspired oxygen fraction is selected (FIO2) and delivered to the patient. This is done automatically via the computer or manually by the doctor. Initially FIO2 is usually that of air (21%) but any other value of FIO2 can be used as the starting point for the experiment. At all times the patient/subject is required to have an arterial oxygen saturation (SpO2) greater than or equal to 0.85. The initial FIO2 may therefore be set to a high level so as to achieve SpO2 ≥ 0.85.

After setting the inspired oxygen level the patients' oxygen system will take time to equilibrate. This usually occurs within 2-5 minutes after the perturbation. The equilibrium of the patients oxygen system is monitored automatically by the "steady state monitor" software in the computer. This functionality substantially reduces the time taken to perform a parameter estimation and is only possible because of the apparatus.

Box D: The assessment of equilibrium can be performed using a number of algorithms, e.g. as follows:

- The arterial oxygen saturation (SpO2) remains constant within a predefined range over a predefined time period.
- 5 2) The difference between the fraction of oxygen in the inspired and expired gas remains constant within a predefined interval over a predefined time period.
 - 3) The calculated oxygen consumption (VO2) remains constant within a predefined interval for a predefined time period.

The oxygen consumption (VO2) is calculated automatically by the computer from the continuously monitored variables using the equation VO2 = f (Vt-Vd) (FIO2- FE'O2) assuming or calculating a value of Vd, or using VO2 = f Vt (FIO2- FEO2), or any variation in this equation where a combination of measurements of end tidal or mixed expired gases are used to estimate the oxygen consumption.

Box E: When equilibrium is achieved a measurement is recorded (Box F).

Box F: This measurement includes the current values of all continuously monitored variables as described previously. It can also include measurements of blood gases in from and arterial or venous blood and a cardiac output measure obtained from equipment e.g. a pulmonary catheter. The last measurements are optional.

Box G: Following a measurement it is decided either automatically by the apparatus or manually by the clinician whether a sufficient number of measurements have been performed, or whether to change the inspired oxygen fraction to a new level and take a further measurement when equilibrium is achieved.

Box H: It is also decided either automatically by the apparatus or manually by the clinician what level of FIO2 should be selected for a new measurement (if necessary). An experiment consists of not less than 2 measurements at varying FIO2 levels, with SpO2 in the range 0.85-1.00. It is important that the setting of FIO2 levels achieve data points with SpO2 well distributed between 0.85-1.00.

Examples of algorithms, which can be used to implement Box G and Box H are included in the next section.

Box I: After an adequate set of measurements has been taken parameters are estimated which describe the patients lung function. Parameter estimation is performed automatically using one or more of the following algorithms:

1) Graphical estimation of displacement(s) of the FIO2/SpO2 curve or the FĒO2/SpO2 curve.

10

Values of inspired or expired oxygen fraction can be plotted against the arterial oxygen saturation (SpO2) and graphical methods used to measure the horizontal (H-shift) and vertical displacement (V-shift) of the data (or interpolated data) from a normal reference range as illustrated in Figure 2.

15

Estimation of the parameters of models of oxygen transport.
 All data collected for each of the measurements can be used with mathematical models of oxygen transport to estimate parameters describing oxygenation.
 Parameters can e.g. be estimated describing the shunting of pulmonary blood (shunt) and either a resistance to oxygen diffusion or a mismatch between the ventilation and perfusion of the lung.

Algorithms for Automating boxes G and H in Fig. 6:

Numerous algorithms can be devised which enable assessment of:

25

- a) Whether a new measurement is required.
- b) What is the target SpO2 for this measurement.
- c) What inspired oxygen fraction (FIO2) setting should be used to obtain the target SpO2

30

These algorithms include those with complete computer automation of points a-c, and where points a-c are assessed using clinical judgement.

Two examples of these algorithms are presented here. The first includes points a and b. The second includes points a and c, using mathematical models of oxygen transport to asses the appropriate FIO2 setting.

5 It should be noted that these algorithms are only illustrations of the control system of ALPE and that any other algorithms which can be used to assess points a, b and c are included in the patent application.

Algorithm 1:This algorithm covers points a and b above, and is illustrated in a flowchart 10 (Fig. 7). It should be noted that if the current FIO2 = 1.0 and the current SpO2 is ≤ 0.85, then the patient is too ill to perform a lung assessment.

Algorithm 2: This algorithm covers points a and c i.e. it assesses whether a measurement is required and estimates the appropriate FIO2 setting for the next measurement given a target SpO2. The algorithm is illustrated in the flowchart Fig. 8. This algorithm uses a mathematical model of oxygen transport with two parameters. Parameters are implemented as stochastic variables and as such have probability distributions as illustrated in Figure 9.

20 In box A (Figure 9) the appropriate a priori estimates are obtained for the parameter distributions. If the patients lung parameters have been investigated previously, or if the patient belongs to a well-defined population there may be well defined a priori distributions for the patient's lung parameters. Alternatively, default parameter settings can be used. An example illustrating probability distributions on a parameter e.g. "shunt" or diffusion resistance "Rdiff" is illustrated in Figure 9.

In box B the predefined target SpO2 level is retrieved from computer storage.

In box C the parameters' probability distributions are updated with the measured data.

This is a Bayesian update of the parameter estimates for the measured values, such that the probability of the parameter values given the measurements

(P(parameters | measurements)) can be calculated from Bayes theorem i.e.

P(parameters | measurements) = P(measurements | parameters) P(parameters)

P(measurements)

The output of this process being revised probability distributions for the patients' lung parameters updated to reflect the new information obtained from the measurements.

These probability distributions are usually somewhat narrower than the *a priori* estimates as illustrated in Fig. 9.

Box D decides whether a further measurement is required. If the updated probability distributions for the patients' lung parameters have a very narrow distribution, then they are estimated with good precision, and no further FIO2 settings or measurements are required. If a further measurement is required then it is necessary to find the appropriate FIO2 setting so as to reach the next target SpO2. This is done in several steps: first the mathematical models are used to predict SpO2 when the FIO2 level is lowered or raised by a predetermined percentage. The predetermined percentage is dependent on the conditions and the patient. SpO2 is then predicted using the updated parameter estimates and the equation:

$$P(SpO2 | (FIO2)) = \sum_{param} P(SpO2 | FIO2, parameters) P(parameters)$$

where P(parameters) is the current joint probability of all the parameter estimates.

The output from this procedure is a set of probability distributions about SpO2 on varying FIO2 values, as illustrated in Figure 10. Next (box F), an FIO2 level is selected. The FIO2 level is chosen such that a small fraction (e.g. 10%) of the predicted probability mass is below the target SpO2 (see Figure 10). Selecting an FIO2 where only a small fraction of the predicted SpO2 probability mass is below the target is a safety feature of this algorithm. Effectively, it means that it is unlikely that the patients SpO2 will fall below the target value on modification of FIO2. After setting the new FIO2 level the SpO2 target is modified and the above procedure repeated.

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CLAIMS

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1. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FEO2,

- 20 PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,
 - the computer being adapted for retrieving and storing at least two measurements being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in
- data storage means associated with the computer, the at least two measurements being conducted at respective levels of oxygen in the gas flow passing into the respiratory system, the computer further being adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on the at least two measurements.
 - 2. A device according to claim 1, wherein the computer further being adapted for determining at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual.

- 3. A device according to claim 1 or 2, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 4. A device according to claim 1 or 3, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising

determining, based on at least two measurements, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection 10 means,

producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

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5. A device according to any of claims 1-4, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously.

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A device according to claim 5, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.

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- 7. A. device according to any of claims 1-6, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item
 5 accordingly if said parameter exceeds said threshold value.
- 8. A device according to any of claim 1-7, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2,
 10 SpO2, PaO2, PpO2) and produce a control data item accordingly.
- 9. A device according to claim 8, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurements.
 - 10. A device according to claim 8, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 11. A device according to any of claims 8-10, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 12. A device according to any of claims 1-11, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.
 - 13. A device according to any of claims 1-12, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 14. A device according to any of the preceding claims, wherein the oxygen saturation in35 the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.

15. A device according to any of claims 1-14, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

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16. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer accordingly, and

second detection means for detecting the level of oxygen (FIO2, FE'O2, FEO2,

PIO2, PEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer being further adapted for performing a procedure at least once, the procedure comprising

determining, based on data stored within the data structure, whether additional measurements are required,

asserting a possible desired target defining a desired output of the first detection means,

producing a possible control data item based on the target, and

retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.

5 17. A device according to claim 16, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, V) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- 15 the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure in data storage means associated with the computer, in which the stored outputs are mutually related and related to the output from the first detection means and the second detection means, and the output from the four detection means can be retrieved simultaneously.
 - 18. A device according to claim 17, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 25 19. A device according to claim 16 or 17, wherein the computer is adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 30 20. A device according to claim 19, wherein at least two respiratory parameters (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.

- 21. A device according to claim 19 or 20, wherein said parameter(s) (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 5 22. A. device according to any of claims 16-21, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.

23. A device according to any of claims 16-22, wherein the computer is adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

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24. A device according to claim 23, wherein the assessment of change in oxygen level in the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).

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- 25. A device according to claim 23, wherein the assessment of change in oxygen level in the inspired gas is based on the rate of change of the output of at least one of the detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 25 26. A device according to any of claims 23-25, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one gas, in response to said control data item relating to the assessed change in oxygen level from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.

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27. A device according to any of claims 16-26, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

accordingly, and

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- 28. A device according to any of claims 16-27, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 29. A device according to any of claims 16-29, wherein the oxygen saturation in the blood 5 circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
 - 30. A device according to any of claims 16-29, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial blood stream.

31. A device for determining one or more respiratory parameters relating to an individual, comprising

a gas flow device having means for conducting a flow of inspiratory gas from an inlet opening to the respiratory system of the individual and a flow of expiratory gas from the respiratory system of the individual to an outlet opening,

a gas-mixing unit for supplying a substantially homogeneous gas to the inlet opening of the gas flow device,

first supply means for supplying a first gas to an inlet of the gas mixing unit and having first control means for controlling the flow of the first gas,

second supply means for supplying a second gas having an oxygen fraction different to the gas supplied from the first supply means to an inlet of the gas mixing unit and having second control means for controlling the flow of the second gas,

a computer for determining said one or more respiratory parameters,

first detection means for detecting the level of oxygen (SaO2, SpO2, PaO2, PpO2) in the blood circulation of the individual and producing an output to the computer

second detection means for detecting the level of oxygen (FIO2, FEO2, FEO2,

PIO2, PE'O2, FEO2) in the gas flow passing into or out of the respiratory system of the individual and producing an output to the computer accordingly,

the computer being adapted for retrieving and storing at least a first measurement being the concurrent output produced by the first detection means and the second detection means within a data structure, in which the two stored outputs are mutually related, in data storage means associated with the computer, the computer further being adapted to asses the appropriate change in oxygen level in the inspired gas (FIO2) from

the current oxygen level (FIO2) so as to achieve a given desired target oxygen level in the blood (SaO2, SpO2, PaO2, PpO2) and produce a control data item accordingly.

- 32. A device according to claim 31, wherein the assessment of change in oxygen level in 5 the inspired gas is based on a predefined set of data representing statistical distributions of parameters stored within data storage means associated with the computer and on said measurement(s).
- 33. A device according to claim 31, wherein the assessment of change in oxygen level in
 the inspired gas is based on the rate of change of the output of at least one of the
 detection means in response to a change in oxygen level (FIO2) in the inspired gas flow.
- 34. A device according to any of claims 31-33, wherein the computer is adapted to operate the control means for controlling the flow to the gas mixing unit of at least one
 15 gas, in response to said control data item from the computer so as to change the oxygen level (FIO2) in the inspired gas flow accordingly.
 - 35. A device according to any of claims 31 to 34, wherein the computer further is adapted for performing a procedure at least once, the procedure comprising
- determining, based on at least one measurement, whether additional measurements are required,
 - asserting a possible desired target defining a desired output of the first detection means.
 - producing a possible control data item based on the target, and retrieving and storing, in the data structure, additional measurement results being the concurrent output produced by the first detection means and the second detection means.
- 36. A device according to any of claims 31-35, wherein the second detection means are arranged for detecting the level (FIO2, PIO2) of oxygen in the gas flow passing into the respiratory system, and the device further comprises

third detection means for detecting the level (FE'O2, FĒO2, PE'O2, PĒO2) of oxygen in the gas flow passing out of the respiratory system and producing an output to the computer accordingly, and

fourth detection means for detecting variables (Vt, f, \dot{V}) of the gas flow passing the respiratory system and producing an output to the computer accordingly, said output being sufficient for the computer to establish the volume flow of gas passing the respiratory system,

- 5 the computer being adapted for retrieving and storing output from the third detection means and the fourth detection means within the data structure relating these stored output mutually as well as with the output from the first detection means and the second detection means retrieved simultaneously.
- 10 37. A device according to claim 36, wherein the computer further being adapted for establishing, based on said measurement(s), the oxygen consumption (VO2) of the individual.
- 38. A device according to any of claims 31-37, wherein the computer is adapted for determining at least one respiratory parameter (Rdiff, shunt, \dot{V}/\dot{Q} , H-shift, V-shift) being descriptive of the condition of the individual, the determination being based on at least two measurements.
- 39. A device according to claim 38, wherein at least two respiratory parameters (Rdiff, 20 shunt, \dot{V}/\dot{Q} , H-shift, V-shift) are determined.
 - 40. A device according to claim 38 or 39, wherein said parameter(s) (Rdiff, shunt, V/\dot{Q} , H-shift, V-shift) is/are generalised parameters being comparable to similar parameter(s) determined for other individuals.
- 41. A. device according to any of claims 31-40, wherein the computer is adapted to determine a parameter relating to an equilibrium state of the overall oxygen uptake or consumption of the individual based on the output of at least one of the detection means, to compare said parameter with a predefined threshold value and to produce a control data item accordingly if said parameter exceeds said threshold value.
 - 42. A device according to any of claims 31-41, wherein one gas is atmospheric air and another gas has an oxygen fraction higher than that of atmospheric air, preferably in the range 0.85 to 1.00.

- 43. A device according to any of claims 31-42, wherein one gas is atmospheric air and another gas has an oxygen fraction in the range of 0.00 to 0.21, preferably 0.00 to 0.05.
- 5 44. A device according to any of claims 31-43, wherein the oxygen saturation in the blood circulation of the individual is in the range of 65 to 100%, preferably 85 to 100%.
- 45. A device according to any of claims 31-43, wherein the first detection means is arranged for detecting a parameter relating to the saturation level of oxygen in the arterial10 blood stream.
 - 46. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is an apparently healthy individual.
 - 47. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is considered to have a risk of suffering from hypoxemia.
- 48. Method for determining one or more respiratory parameters by means of a device according to any of the preceding claims, wherein the individual is suffering from hypoxemia.
- 49. Method according to claim 48, wherein the individual is suffering from one or more disease(s) selected from the group(s) comprising left sided heart failure, adult respiratory distress syndrome, pneumonia, postoperative hypoxemia, pulmonary fibrosis, toxic pulmonary lymphoedema, pulmonary embolisms, chronic obstructive pulmonary disease and cardiac shunting.
- 30 50. A computer system comprising at least one general purpose computer having one or more computer programs stored within data storage means associated therewith, the computer system being arranged for as well as being adapted for determining one or more respiratory parameters according to any of claims 1-49.

51. A computer program product being adapted to enable a computer system comprising at least one general purpose computer having data storage means associated therewith and being arranged suitably to determine one or more respiratory parameters according to any of claims 1-49.

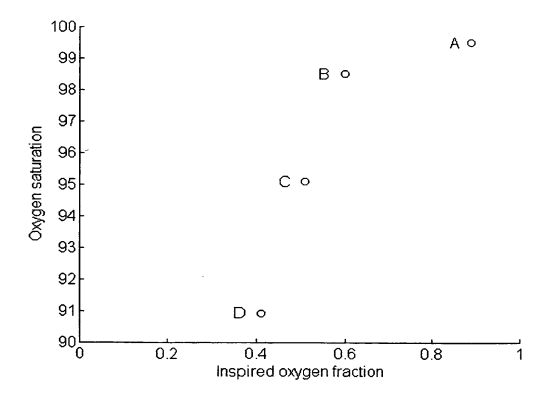


Fig. 1

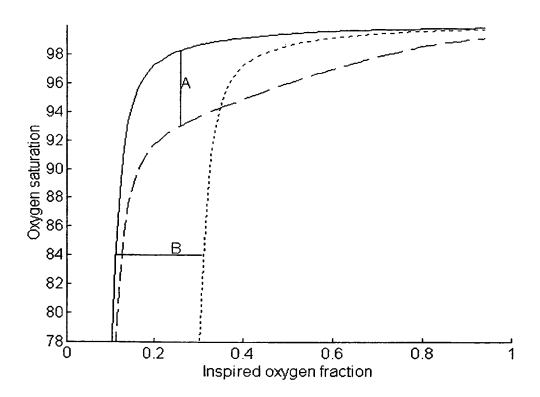


Fig. 2

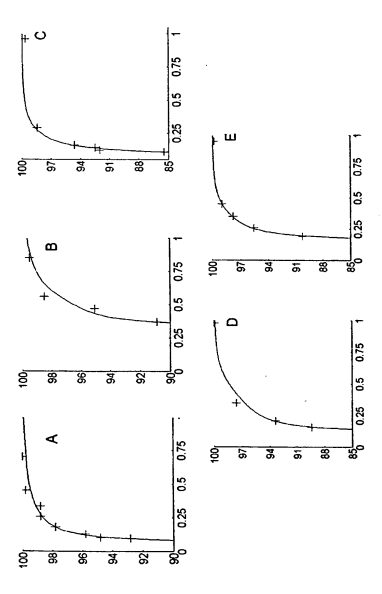


Fig. 3

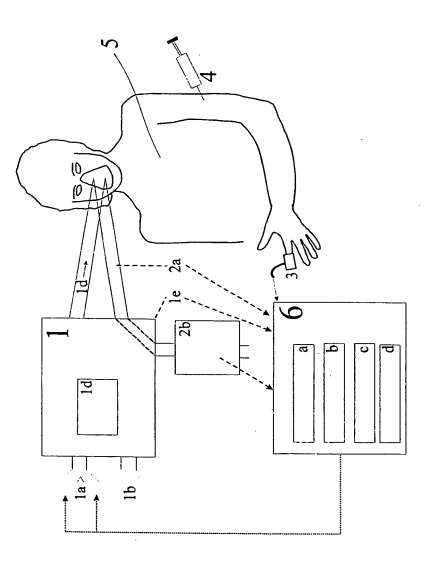


Fig. 4

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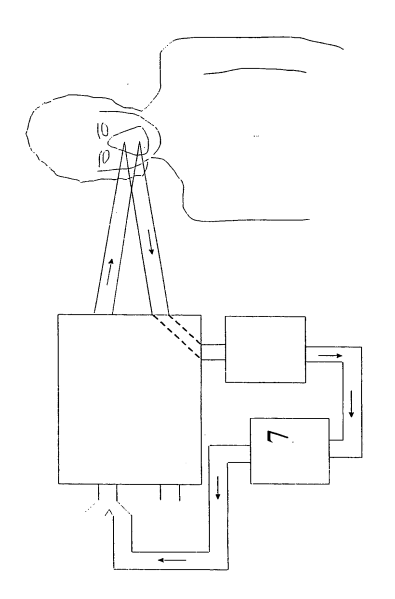


Fig. 5

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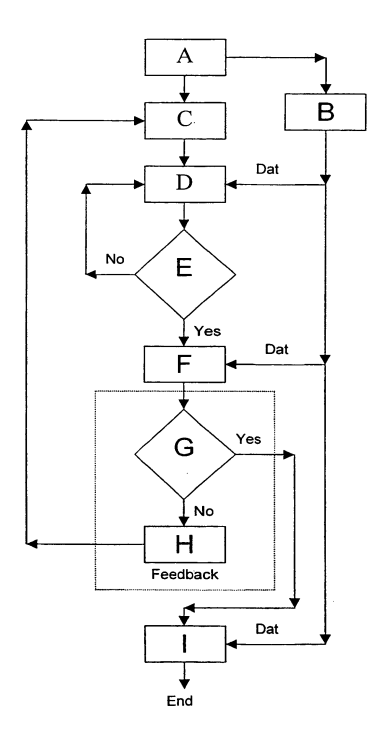


Fig. 6

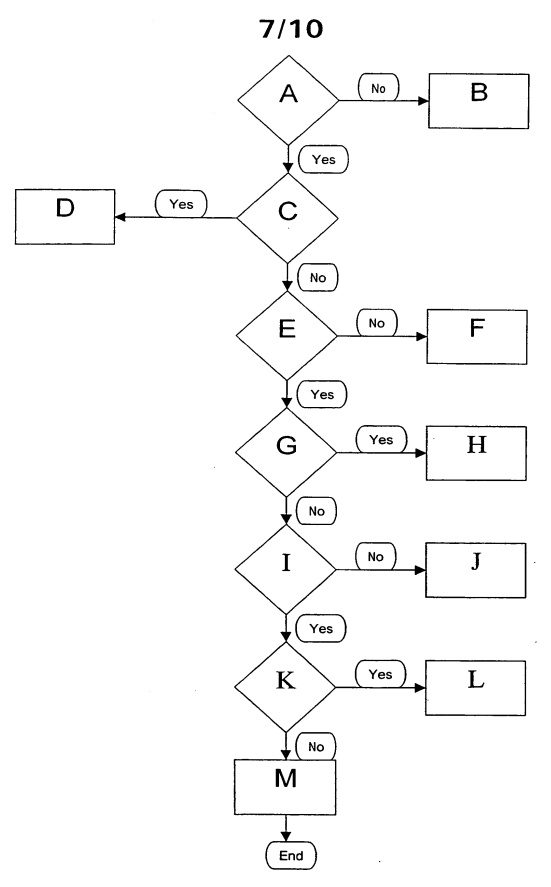


Fig. 7

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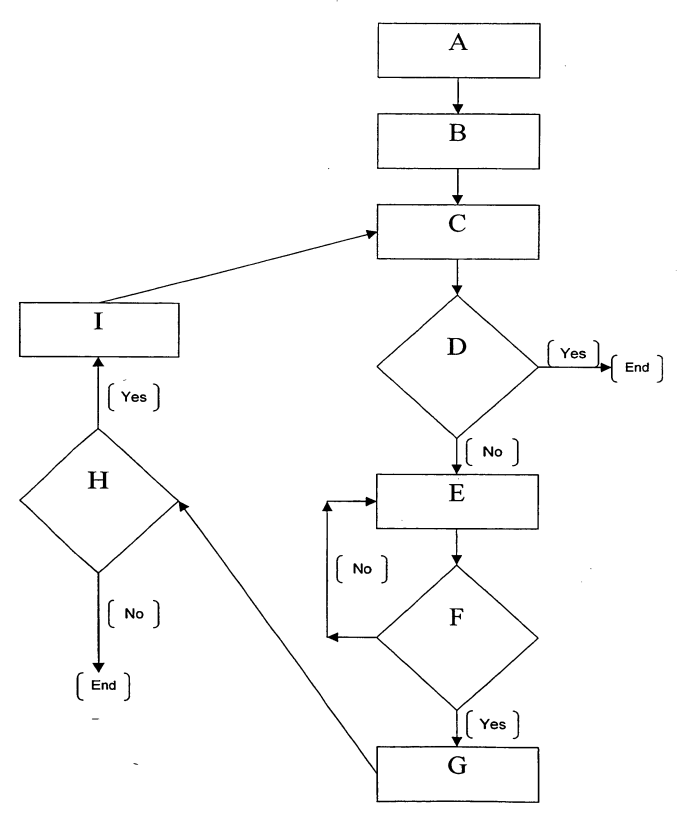


Fig. 8

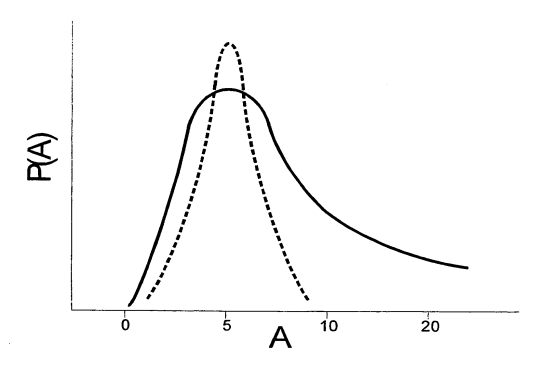


Fig. 9

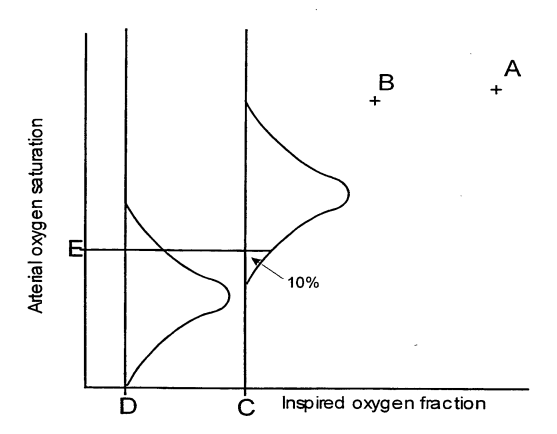


Fig. 10

CLASSIFICATION OF SUBJECT MATTER IPC7: A61B 5/08, A61M 16/00 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) IPC7: A61B, A61M Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. Category* EP 0753320 A1 (B. LACHMANN), 15 January 1997 1-3,31 X (15.01.97), column 7, line 41 - column 9, line 2, abstract 4-30,32-51 column 7, line 41 - column 9, line 2, A abstract US 5103814 A (T. MAHER), 14 April 1992 (14.04.92), 1-51 column 3, line 3 - line 55 1-51 EP 0502270 A1 (HAMAMATSU PHOTONICS K.K.), A 9 Sept 1992 (09.09.92), page 4, line 46 - page 5, line 4, abstract See patent family annex. Further documents are listed in the continuation of Box C. later document published after the international filing date or priority Special categories of cited documents: date and not in conflict with the application but cited to understand document defining the general state of the art which is not considered the principle or theory underlying the invention to be of particular relevance "X" document of particular relevance: the claimed invention cannot be erlier document but published on or after the international filing date considered novel or cannot be considered to involve an inventive document which may throw doubts on priority claim(s) or which is step when the document is taken alone cited to establish the publication date of another citation or other special reason (as specified) document of particular relevance: the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination document referring to an oral disclosure, use, exhibition or other being obvious to a person skilled in the art document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search **27** 06, 2000 <u>25 May 2000</u> Authorized officer Name and mailing address of the International Searching Authority European Patent Office P.B. 5818 Patentiaan 2 Ulrika Andersson/AE MI -2280 HV Rimmilk Tel(+31-70)340-2040. Tx 31 651 epo nl. Telephone No.

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